nature portfolio

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Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our <u>Editorial Policies</u> and the <u>Editorial Policy Checklist</u>.

Please do not complete any field with "not applicable" or n/a. Refer to the help text for what text to use if an item is not relevant to your study. For final submission: please carefully check your responses for accuracy; you will not be able to make changes later.

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For	all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.	
n/a	Confirmed	
	\mathbf{Q} The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement	
\bigvee	A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly	
\bigvee	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.	
\bigvee	A description of all covariates tested	
	🗹 A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons	
	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)	
abla	For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted Give <i>P</i> values as exact values whenever suitable.	
abla	For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings	
abla	For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes	
\bigvee	\square Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated	
,	Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.	
Sof	ftware and code	
Policy information about <u>availability of computer code</u>		

Data collection

MIDAS data from IQVIA was presented in excel format

Data analysis

Statistical Analysis System (SAS) v9.4 and R Foundation for Statistical Computing version 3.6.0 are used for data analysis. R codes adopted in this study have been made available in GitHub repository at (https://github.com/adrienneylc/Gabapentinoids).

For manuscripts utilizing

reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

The MIDAS data from IQVIA are available under restricted access for licensing reasons, access can be obtained by entering into additional licensing agreement with IQVIA. The raw MIDAS data are protected and are not publicly available due to data protection agreement with IQVIA. With additional data use agreement and permission from IQVIA, MIDAS data will be made available from the corresponding authors upon request. Source data of tables and figures presented are provided with this paper.

Research invo	lving hur	man participants, their data, or biological material
Policy information aboand sexual orientation		ith <u>human participants or human data</u> . See also policy information about <u>sex, gender (identity/presentation), hnicity and racism</u> .
Reporting on sex an	ıd gender	No Applicable
Reporting on race, other socially releva	• • •	No Applicable
Population characte	eristics	No Applicable
Recruitment		No Applicable
Ethics oversight		No Applicable
Note that full information	n on the appro	oval of the study protocol must also be provided in the manuscript.
Field-spec	ific re	porting
Please select the one	below that is	the best fit for your research. If you are not sure, read the appropriate sections before making your selection.
Life sciences	□ Ве	ehavioural & social sciences
		Ill sections, see nature.com/documents/nr-reporting-summary-flat.pdf
.,		
Life science	ces stu	ıdy design
All studies must disclo	ose on these i	points even when the disclosure is negative.
Figu Sample size dose	ire 4 presents the av	erage annual percentage change of gabapentinoid consumption over the span of 11 years, from 2008 to 2018. Gabapentinoid consumption is presented as defined dail shabitants per day. The mid-year population estimates of each country was obtained from the UN Population Division in 2019 and each individual country has a different
Data exclusions	No data was exclud	led in the main analysis of the study.
Replication T		have conducted the statistical analysis independently using the same set of MIDAS data from IQVIA. In the manuscript were cross-checked by the first two authors using different statistical analysis programmes (SAS and R) to ensure reproducibility. All attempts at essful.
Randomization Ou	Our longitudinal trend study utilise real-world sales data to describe the consumption trend of gabapentinoids and randomisation is not applicable.	
Blinding	Our longitudinal tr	end study utilise real-world sales data to describe the consumption trend of gabapentinoids and blinding is not applicable.
Behaviour	al & s	ocial sciences study design
All studies must disclo	ose on these p	points even when the disclosure is negative.
Study description		
Research sample		
Sampling strategy		
Data collection		

Timing Data exclusions

Non-participation

Randomization

Ecological, e	volutionary	& environmental sciences study design
All studies must disclose on	these points even when	the disclosure is negative.
Study description		
Research sample		
Sampling strategy		
Data collection		
Timing and spatial scale		
Data exclusions		
Reproducibility		
Randomization		
Blinding		
Did the study involve field	d work? Yes] No
Field work, collect	tion and transpo	rt
Field conditions		
Location		
Access & import/export		
Disturbance		
		aterials, systems and methods materials, experimental systems and methods used in many studies. Here, indicate whether each material,
system or method listed is rele	vant to your study. If you are	e not sure if a list item applies to your research, read the appropriate section before selecting a response.
Materials & experime	ntal systems	Methods
n/a Involved in the study Antibodies		n/a Involved in the study
Antibodies Likaryotic cell lines		✓ ChIP-seq ✓ Flow cytometry
Palaeontology and archaeology		MRI-based neuroimaging
Animals and other o	rganisms	
Clinical data Dual use research of	f concern	
Plants	COMMENT	

Antibodies

Antibodies used	
Validation	

Eukaryotic celi line	28
Policy information about <u>ce</u>	Il lines and Sex and Gender in Research
Cell line source(s)	
Authentication	
Mycoplasma contamination	on
Commonly misidentified li (See <u>ICLAC</u> register)	ines
Palaeontology and	d Archaeology
Specimen provenance	
Specimen deposition	
Dating methods	
Tick this box to confirm	n that the raw and calibrated dates are available in the paper or in Supplementary Information.
Ethics oversight	
Note that full information on th	e approval of the study protocol must also be provided in the manuscript.
Animals and other	r research organisms
Policy information about stu Research	udies involving animals; ARRIVE guidelines recommended for reporting animal research, and Sex and Gender in
Laboratory animals	
Wild animals	
Reporting on sex	
Field-collected samples	
Ethics oversight	
Note that full information on th	ne approval of the study protocol must also be provided in the manuscript.
Clinical data	
Policy information about <u>cli</u> All manuscripts should comply v	nical studies with the ICMJE guidelines for publication of clinical research and a completed CONSORT checklist must be included with all submissions.
Clinical trial registration	
Study protocol	
Data collection	
Outcomes	

Dual use research of concern

Policy information about <u>dual use research of concern</u>

Hazards

Could the accidental, deliberate or reckless misuse of agents or technologies generated in the work, or the application of information presented in the manuscript, pose a threat to:

No Yes			
Public health			
National security			
Crops and/or livestocl	Crops and/or livestock		
Ecosystems			
Any other significant a	area		
Experiments of concern			
Does the work involve any o	of these experiments of concern:		
No Yes			
Demonstrate how to	render a vaccine ineffective		
Confer resistance to t	herapeutically useful antibiotics or antiviral agents		
Enhance the virulence	e of a pathogen or render a nonpathogen virulent		
Increase transmissibil	ity of a pathogen		
Alter the host range o	of a pathogen		
Enable evasion of diag	gnostic/detection modalities		
	ation of a biological agent or toxin		
Any other potentially	harmful combination of experiments and agents		
Plants			
Seed stocks			
Novel plant genotypes			
Authentication			
ChIP-seq			
Data deposition			
	nd final processed data have been deposited in a public database such as <u>GEO</u> .		
Confirm that you have d	eposited or provided access to graph files (e.g. BED files) for the called peaks.		
Data access links			
May remain private before publicati	ion.		
Files in database submission	n (
Genome browser session (e.g. <u>UCSC</u>)			
Methodology			
Replicates			
Sequencing depth			
Antibodies			
Peak calling parameters			
Data quality			
Software			

Flow Cytometry	
The axis scales are clearly visib	er and fluorochrome used (e.g. CD4-FITC). ole. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers). h outliers or pseudocolor plots. of cells or percentage (with statistics) is provided.
Methodology	
Sample preparation	
Instrument	
Software	
Cell population abundance	
Gating strategy	
Tick this box to confirm that a	figure exemplifying the gating strategy is provided in the Supplementary Information.
Magnetic resonance in	naging
	<u>laging</u>
Experimental design Design type	
Design type Design specifications	
Behavioral performance measure	
bellavioral performance measure	
Imaging type(s)	
Field strength	
Sequence & imaging parameters	
Area of acquisition	
Diffusion MRI Used	☐ Not used
Preprocessing	
Preprocessing software	
Normalization	
Normalization template	
Noise and artifact removal	
Volume censoring	
Statistical modeling & inferer	nce
Model type and settings	
Effect(s) tested	
Specify type of analysis: Wh	oole brain ROI-based Both

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Statistic type for inference		
(See Eklund et al. 2016)		
Correction		
Models & analysis		
n/a Involved in the study		
Functional and/or effective co	onnectivity	
Graph analysis		
Multivariate modeling or pred	lictive analysis	
Functional and/or effective connect	tivity	
Graph analysis		

Multivariate modeling and predictive analysis